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WHAT IS CLAIMED IS:

- 1. A method for improving an electron density map representing a crystal structure comprising:
- (a) obtaining by x-ray diffraction observed structure factor amplitudes for a plurality of reflections from the crystal structure;
- (b) selecting a starting set of crystallographic phases to combine with the observed structure factor amplitudes to form a first set of structure factors;
- (c) deriving a first electron density map from the first set of structure factors;
- (d) identifying features of the first electron density map to obtain expected distributions of electron density;
- (e) making a comparison between the first electron density map and the expected distribution of electron density;
- (f) estimating how changes in the crystallographic phase of a reflection *k* affect the comparison;
- (g) establishing crystallographic phase probability distributions from the comparisons for the possible crystallographic phases of reflection k;
- (h) repeating steps (c) through (g) as *k* is indexed through all of the plurality of reflections;
- (i) deriving an updated electron density map using crystallographic phases determined to be most probable from the crystallographic phase probability distributions for each one of the reflections; and
- (j) repeating steps (d) through (i) to obtain a final set of crystallographic phases with minimum bias from known electron density maps.
- 2. The method of Claim 1, wherein identifying features of the electron density map includes making probability estimates of whether each point in the map is located in a solvent region or a crystal structure region.
- 3. The method of Claim 1, wherein identifying features of the election density map includes estimates of whether the electron density at each point in the map is related by non-crystallographic symmetry to electron density at another point in the map.

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- 4. The method of Claim 1, includes estimates of whether a structural motif is located at each point in the map.
 - 5. The method of Claim 4, wherein the structural motif is a helix.
- 6. The method of any one of Claims 1, 2, 3, or 4, wherein the crystallographic phase probability distributions are log-likelihood functions.
- 7. The method of Claim 1, further including the steps of calculating first and second derivatives for the crystallographic phase probability distributions with respect to the structure factors; and

applying an FFT-based algorithm to determine the most probable crystallographic phase probability distributions.

8. The method of Claim 1, wherein the step of selecting a starting set of crystallographic phases includes;

selecting a model crystal structure having similarities to the crystal structure being examined;

assigning a low weighting factor to structure factors of the model crystal structure; and

combining the weighted structure factors with the observed structure factors for deriving the first electron density map.